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- (71) Applicant (for all designated States except US): SOLVAY PHARMACEUTICALS B.V. [NL/NL]; C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): KRUSE, Cornelis, G. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL). LANGE, Josephus, H.M. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL). HERREMANS, Arnoldus, H.J. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL). VAN STUIVENBERG, Herman, H. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL).

- (74) Agent: MUIS, Maarten; OCTROOIBUREAU ZOAN B.V., P.O. Box 140, NL-1380 AC Weesp (NL).
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$$\begin{array}{c|c}
 & O \\
 & N \\
 & R_3
\end{array}$$

$$\begin{array}{c|c}
 & R_2 \\
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$$\begin{array}{c|c}
 & R_3
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(54) Title: 1H-IMIDAZOLE DERIVATIVES HAVING CB₁ AGONISTIC, CB₁ PARTIAL AGONISTIC OR CB₁- ANTAGONISTIC ACTIVITY

(57) Abstract: AbstractThe present invention relates to a group of novel 1H-imidazole derivatives, to methods for the preparation of these compounds, and to pharmaceutical compositions containing one or more of these compounds as an active component. These 1H-imidazole derivatives are potent cannabinoid-CB1 receptor agonists, partial agonists or antagonists, useful for the treatment of psychiatric and neurological disorders, as well as and other diseases involving cannabinoid neurotransmission. The compounds have the general formula (I) wherein R and R1-R4 have the meanings given in the specification.



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1H-Imidazole derivatives having CB₁ agonistic, CB₁ partial agonistic or CB₁-antagonistic activity

The present invention relates to a group of novel 1H-imidazole derivatives, to methods for the preparation of these compounds, and to pharmaceutical compositions containing one or more of these compounds as an active component.

These 1H-imidazole derivatives are potent cannabinoid-CB₁ receptor agonists, partial

agonists or antagonists, useful for the treatment of psychiatric and neurological disorders, as well as and other diseases involving cannabinoid neurotransmission.

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Cannabinoids are present in the Indian hemp Cannabis sativa and have been used as medicinal agents for centuries (Mechoulam, R. and Feigenbaum, J. J. Prog. Med. Chem. 1987, 24, 159). However, only within the past ten years the research in the cannabinoid area has revealed pivotal information on cannabinoid receptors and their (endogenous) agonists and antagonists. The discovery and the subsequent cloning of two different subtypes of cannabinoid receptors (CB₁ and CB₂) stimulated the search for novel cannabinoid receptor antagonists (Munro, S. et al., Nature 1993, 365, 61. Matsuda, L. A. and Bonner, T. I. Cannabinoid Receptors, Pertwee, R. G. Ed. 1995, 117, Academic Press, London). In addition, pharmaceutical companies became interested in the development of cannabinoid drugs for the treatment of diseases connected with disorders of the cannabinoid system (Consroe, P. Neurobiology of Disease 1998, 5, 534. Pop, E. Curr. Opin. In CPNS Investigational Drugs 1999, 1, 587. Greenberg, D. A. Drug News Perspect. 1999, 12, 458. Pertwee, R.G., Progress in Neurobiology 2001, 63, 569). Hitherto, several CB₁ receptor antagonists are known. Sanofi disclosed their diarylpyrazole congeners as selective CB₁ receptor antagonists. A representative example is SR-141716A (Dutta, A.K. et al., Med. Chem. Res. 1994, 5, 54. Lan, R. et al., J. Med. Chem. 1999, 42, 769. Nakamura-Palacios, E. M. et al., CNS Drug Rev. 1999, 5, 43). CP-272871 is a pyrazole derivative, like SR141716A, but less potent and less CB₁ receptor subtypeselective than SR141716A (Meschler, J. P. et al., Biochem. Pharmacol. 2000, 60, 1315). Aminoalkylindoles have been dis-closed as CB₁ receptor antagonists. A representative example is lodopravadoline (AM-630), which was introduced in 1995. AM-630 is a moderately active CB₁ receptor antagonist, in some assays behaving as a weak partial agonist (Hosohata, K. et al., Life Sc. 1997, 61, PL115). Researchers from Eli Lilly described aryl-aroyl substituted benzofurans as selective CB1 receptor antagonists (e.g. LY-320135) (Felder, C. C. et al., J. Pharmacol. Exp. Ther. 1998, 284, 291). 3-Alkyl-5,5'-diphenylimidazolidine-diones were described as cannabinoid receptor ligands, which were indicated to be cannabinoid antagonists (Kanyonyo, M. et al., Biorg. Med.Chem. Lett. 1999, 9, 2233). Aventis Pharma claimed diarylmethyleneazetidine analogs as CB₁ receptor antagonists (Mignani, S. et al., Patent FR 2783246, 2000; Chem. Abstr. 2000, 132, 236982). Tricyclic pyrazoles were claimed by Sanofi-Synthelabo as CB1 antagonists (Barth, F. et al. Patent WO 0132663, 2001; Chem. Abstr. 2001, 134, 340504). Interestingly, many CB₁ receptor

antagonists have been reported to behave as inverse agonists *in vitro* (Landsman, R. S. *et al.*, *Eur. J. Pharmacol.* **1997**, *334*, R1). Pyrazole cannabinoids have also been reported as CB₁ receptor partial agonists showing *in vivo* cannabimimetic effects (Wiley, J. L. *et al.*, *J. Pharmacol. Exp. Ther.* **2001**, *296*, 1013). A number of classes of CB₁ receptor agonists are known such as for example the classical cannabinoids (*e.g.* Δ⁹-THC), non-classical cannabinoids, aminoalkylindoles and eicosanoids (*e.g.* anandamide). Reviews provide a nice overview of the cannabinoid research area (Mechoulam, R. *et al.*, *Prog. Med. Chem.* **1998**, *35*, 199. Lambert, D. M. *Curr. Med. Chem.* **1999**, *6*, 635. Mechoulam, R. *et al.*, *Eur. J. Pharmacol.* **1998**, *359*, 1. Williamson, E. M. and Evans, F. J. *Drugs* **2000**, *60*, 1303. Pertwee, R. G. *Addiction Biology* **2000**, *5*, 37. Robson, P. *Br. J. Psychiatry* **2001**, *178*, 107. Pertwee, R. G. *Prog. Neurobiol.* **2001**, *63*, 569. Goya, P. and Jagerovic, N. *Exp. Opin. Ther. Patents* **2000**, *10*, 1529. Pertwee, R. G. *Gut* **2001**, *48*, 859).

15 It has now surprisingly been found that the novel 1H-imidazole derivatives of the formula (I), prodrugs thereof and salts thereof, are potent agonists, partial agonists or antagonists on cannabinoid-CB₁ receptors

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wherein

- R represents phenyl, thienyl, 2-pyridinyl, 3-pyridinyl, 4-pyridinyl, pyriazinyl, pyridazinyl or triazinyl, which groups may be substituted with 1, 2, 3 or 4 substituents Y, which can be the same or different, from the group C₁₋₃-alkyl or alkoxy, hydroxy, halogen, trifluoromethyl, trifluoromethylthio, trifluoromethoxy, nitro, amino, mono- or dialkyl (C₁₋₂)-amino, mono- or dialkyl (C₁₋₂)-amido, (C₁₋₃)-alkoxycarbonyl, carboxyl, cyano, carbamoyl and acetyl, or R represents naphtyl, with the proviso that when R is 4-pyridinyl, R₄ represents a halogen atom or a cyano, carbamoyl, formyl, acetyl, trifluoroacetyl, fluoroacetyl, propionyl, sulfamoyl, methanesulfonyl, methylsulfanyl or branched or unbranched C₁₋₄ alkyl group, which C₁₋₄ alkyl group may be substituted with 1-3 fluoro atoms or with a bromo, chloro, iodo, cyano or hydroxy group,
- R₁ represents phenyl or pyridinyl, which groups may be substituted with 1-4 substituents Y, which can be the same or different, wherein Y has the above mentioned meaning, or R₁ represents pyrimidinyl, pyrazinyl, pyridazinyl or triazinyl, which groups may be substituted with 1-2 substituents Y, which can be the same or different or R₁ represents a five-membered aromatic heterocyclic ring having one or two heteroatoms from the group (N, O, S), which heteroatoms can

be the same or different, which five-membered aromatic heterocyclic ring may be substituted with 1-2 substituents Y, which can be the same or different or R₁ represents naphtyl,

R₂ represents H, branched or unbranched C₁₋₈ alkyl, C₃₋₈ cycloalkyl, C₃₋₈ alkenyl,
 C₅₋₈ cycloalkenyl which groups may contain a sulfur, oxygen or nitrogen atom,

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- R₃ represents branched or unbranched C₂₋₈ alkyl, C₁₋₈ alkoxy, C₅₋₈ cycloalkyloxy, C₃₋₈ cycloalkyl, C₅₋₁₀ bicycloalkyl, C₆₋₁₀ tricycloalkyl, C₃₋₈ alkenyl, C₅₋₈ cycloalkenyl, which groups may optionally contain one or more heteroatoms from the group (O, N, S) and which groups may be substituted with a hydroxy group or 1-2 C₁₋₃ alkyl groups or 1-3 fluoro atoms, or R₃ represents a benzyl or phenethyl group which aromatic rings may be substituted with 1-5 substituents Z, which can be the same or different, from the group C₁₋₃-alkyl or alkoxy, hydroxy, halogen, trifluoromethyl, trifluoromethylthio, trifluoromethoxy, nitro, amino, mono- or dialkyl (C₁₋₂)-amino, mono- or dialkyl (C₁₋₂)-amido, (C₁₋₃)-alkylsulfonyl, dimethyl-sulfamido, C₁₋₃-alkoxycarbonyl, carboxyl, trifluoromethylsulfonyl, cyano, carbamoyl, sulfamoyl and acetyl, or R₃ represents a phenyl or pyridinyl group, which groups are substituted with 1-4 substituents Z, wherein Z has the meaning as indicated above.
- or R₃ represents a pyridinyl group, or R₃ represents a phenyl group, with the proviso that R₄ represents a halogen atom or a cyano, carbamoyl, formyl, acetyl, trifluoroacetyl, fluoroacetyl, propionyl, sulfamoyl, methanesulfonyl, methylsulfanyl or C₁₄ alkyl group, which C₁₄ alkyl group may be substituted with 1-3 fluoro atoms or with a bromo, chloro, iodo, cyano or hydroxy group,
- or R_3 represents a group NR_5R_6 with the proviso that R_2 represents a hydrogen atom or a methyl group, wherein
 - R₅ and R₆ are the same or different and represent branched or unbranched C₁₋₄ alkyl, or R₅ and R₆ together with the nitrogen atom to which they are bonded form a saturated or unsaturated, monocyclic or bicyclic heterocyclic group having 4 to 10 ring atoms which heterocyclic group contains one or two heteroatoms from the group (N, O, S), which heteroatoms can be the same or different, which heterocyclic group may be substituted with a C₁₋₃ alkyl group or a hydroxy group, or R₂ and R₃ together with the nitrogen atom to which they are bonded form a saturated or unsaturated heterocyclic group having 4 to 10 ring atoms which heterocyclic group contains one or two heteroatoms from the group (N, O, S), which heteroatoms can be the same or different, which heterocyclic group may be substituted with a C₁₋₃ alkyl group or a hydroxy group,
 - R₄ represents a hydrogen or halogen atom or a cyano, carbamoyl, formyl, acetyl, trifluoroacetyl, fluoroacetyl, propionyl, sulfamoyl, methanesulfonyl, methylsulfanyl or branched or unbranched C₁₄ alkyl group, which C₁₄ alkyl group may be substituted with 1-3 fluoro atoms or with a bromo, chloro, iodo, cyano or a hydroxy group,

Due to the potent CB₁ agonistic, partial agonistic or antagonistic activity the compounds according to the invention are suitable for use in the treatment of psychiatric disorders such as psychosis, anxiety, depression, attention deficits, memory disorders, cognitive disorders, appetite disorders, obesity, addiction, 5 'appetence, drug dependence and neurological disorders such as neurodegenerative disorders, dementia, dystonia, muscle spasticity, tremor, epilepsy, multiple sclerosis, traumatic brain injury, stroke, Parkinson's disease, Alzheimer's disease, epilepsy, Huntington's disease, Tourette's syndrome, cerebral ischaemia, cerebral apoplexy, craniocerebral trauma, stroke, spinal cord injury, neuroinflammatory disorders, plague sclerosis, viral encephalitis, demyelinisation related disorders, as well as for the treatment of pain disorders, including neuropathic pain disorders, and other diseases involving cannabinoid neurotransmission, including the treatment of septic shock, glaucoma, diabetes, cancer, emesis, nausea, gastrointestinal disorders, gastric ulcers, diarrhoea and cardiovascular disorders.

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The affinity of the compounds of the invention for cannabinoid CB₁ receptors was determined using membrane preparations of Chinese hamster ovary (CHO) cells in which the human cannabinoid CB₁ receptor is stably transfected in conjunction with [3H]CP-55,940 as radioligand. After incubation of a freshly prepared cell membrane preparation with the [3H]-ligand, with or without addition of compounds of the invention, separation of bound and free ligand was performed by filtration over glassfiber filters. Radioactivity on the filter was measured by liquid scintillation counting.

25 The cannabinoid CB₁ antagonistic activity of compounds of the invention was determined by functional studies using CHO cells in which human cannabinoid CB₁ receptors are stably expressed. Adenylyl cyclase was stimulated using forskolin and measured by quantifying the amount of accumulated cyclic AMP. Concomitant activation of CB₁ receptors by CB₁ receptor agonists (e.g. CP-55,940 or (R)-WIN-30 55,212-2) can attenuate the forskolin-induced accumulation of cAMP in a concentration-dependent manner. This CB₁ receptor-mediated response can be antagonized by CB₁ receptor antagonists such as the compounds of the invention.

Cannabinoid agonistic of partial agonistic activity of compounds of the invention can be determined according to published methods, such as assessment of in vivo cannabimimetic effects (Wiley, J. L. et al., J. Pharmacol. Exp. Ther. 2001, 296, 1013).

The invention relates both to racemates, mixtures of diastereomers and the individual stereoisomers of the compounds having formula (1).

The compounds of the invention can be brought into forms suitable for administration by means of usual processes using auxiliary substances and/or liquid or solid carrier materials.

5 Suitable synthetic routes for the compounds of the invention are the following:

Synthetic route A

Step 1: ester hydrolysis of a compound having formula (II) wherein R₇ represents a branched or unbranched alkyl group (C₁₋₄) or benzyl group

$$R_1$$
 R_4
 R_4
 R_4
 R_4
 R_4
 R_4

This reaction gives a compound having formula (III)

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wherein R, R_1 and R_4 have the meanings as described above.

Intermediates having formula (II), wherein R_7 represents a branched or unbranched alkyl group (C_{1-4}) or benzyl group can be obtained according to methods known, for example:

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- a) I. K. Khanna et al., J. Med. Chem. 2000, 43, 3168-3185
- b) N. Kudo et al., Chem. Pharm. Bull. 1999, 47, 857-868
- c) K. Tsuji et al., Chem. Pharm. Bull. 1997, 45, 987-995
- d) I. K. Khanna et al., J. Med. Chem. 1997, 40, 1634-1647
- 25 e) M. Guillemet et al., Tetrahedron Lett. 1995, 36, 547-548

<u>Step 2</u>: reaction of a compound having formula (III) with a compound having formula R_2R_3NH wherein R_2 and R_3 have the meanings as described above *via* activating and coupling methods such as formation of an active ester, or in the presence of a coupling reagent such as DCC, HBTU, BOP or similar reagents. This reaction gives a desired 1H-imidazole derivative having formula (I). (For more information on activating and coupling methods see: M. Bodanszky and A. Bodanszky: The Practice of Peptide Synthesis, Springer-Verlag, New York, 1994; ISBN: 0-387-57505-7).

Alternatively, a compound having formula (III) is reacted with a halogenating agent, for example thionyl chloride (SOCl₂). This reaction gives the corresponding carbonyl chloride (IV).

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$$R_1$$
 R_4
 R
 R
 R
 R
 R

Reaction of a compound having formula (IV) with a compound having formula R_2R_3NH wherein R_2 and R_3 have the meanings as described above, yields a 1H-imidazole derivative having formula (I). This reaction is preferably carried out in the presence of an organic base such as for example diisopropylethylamine (DIPEA) or triethylamine.

Alternatively, a compound having formula (II) is reacted in an amidation reaction with a compound having formula R₂R₃NH wherein R₂ and R₃ have the meanings as described above to give a 1H-imidazole derivative having formula (I).

Synthetic route B

Reaction of a compound having formula (II), wherein R₄ represents hydrogen and wherein R, R₁ and R₇ have the meanings as described above for compound (II), with a compound having general formula R₄'-X, wherein X represents a leaving group and R₄' represents a C₁₋₄ alkyl group, which alkyl group may be substituted with 1-3 fluoro atoms or wherein R₄' represents a cyano, formyl, acetyl, trifluoroacetyl, fluoroacetyl, methylsulfanyl or propionyl moiety, or a halogen atom. This reaction is carried out in the presence of a strong non-nucleophilic base such as lithium diisopropylamide (LDA), preferably under anhydrous conditions in an inert organic solvent, for example tetrahydrofuran, and yields a compound having formula (II)

$$R_{1} \xrightarrow{N} R_{4}$$

$$R$$

$$R$$

$$R$$

$$R$$

$$R$$

$$R$$

$$R$$

$$R$$

$$R$$

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wherein R, R_1 and R_7 have the meanings as described hereinabove and R_4 represents a C_{1-4} alkyl group, which alkyl group may be substituted with 1-3 fluoro atoms or wherein R_4 represents a cyano, formyl, acetyl, trifluoroacetyl, fluoroacetyl, methylsulfanyl or propionyl group, or a halogen atom.

Compounds of general formula (II) which have been obtained according to synthesis route B can be converted to compounds of general formula (I) analogously to the procedures described in synthesis route A, step 1 of route A or step 2 of route A (see above).

Synthetic route C

Compounds having formula (II)

$$R_1 \xrightarrow{N} R_4$$
(II)

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wherein R_4 represents a branched or unbranched C_{1-4} alkyl group, which C_{1-4} alkyl group may be substituted with 1-3 fluoro substituents and wherein R, R_1 have the meanings given above and R_7 represents a branched or unbranched alkyl group (C_{1-4}) or benzyl group can be synthesized by reacting a compound having formula (V) or its tautomer

wherein R and R₁ have the meanings given above, with a compound having formula (VI)

wherein R₄ represents a branched or unbranched C₁₋₄ alkyl group, which C₁₋₄ alkyl group may be substituted with 1-3 fluoro atoms and R₈ represents a leaving group, for example a bromo substituent, and R₇ represents a branched or unbranched alkyl group (C₁₋₄) or benzyl group. The reaction is preferably carried out in an organic solvent, for example in 2-propanol or in N-methyl-2-pyrrolidinone (NMP). The addition of an acid like trifluoroacetic acid (TFA) during the reaction may enhance the formation of the compounds having formula (II).

(For more information on the leaving group concept see: M. B. Smith and J. March: Advanced organic chemistry, p. 275, 5th ed., (2001) John Wiley & Sons, New York, ISBN: 0-471-58589-0).

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Compounds of general formula (II) which have been obtained according to synthesis route C can be converted to compounds of general formula (I) analogously to the

procedures described in synthesis route A, step 1 of route A or step 2 of route A (see above).

Compounds of the invention having formula (VI) can be obtained according to methods known, for example: P. Seifert et al., Helv. Chim. Acta, 1950, 33, 725.

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Synthetic route D

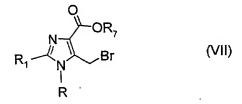
Reaction of a compound having formula (II)

$$R_1$$
 R_4
 R
 R
 R
 R
 R
 R
 R
 R
 R

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wherein R_4 represents a methyl group and R, R_1 have the meanings given above and R_7 represents a branched or unbranched alkyl group (C_{1-4}) or benzyl group with a regioselective brominating compound such as N-bromo-succinimide (NBS) in an organic solvent such as CCl_4 in the presence of a free-radical initiator like dibenzoyl peroxide gives a compound of formula (VII)



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wherein R, R₁ and R₇ have the meanings given above. Reaction of a compound having formula (VII) (analogous to the method described in Mathews, W.B. et al., J. Label. Compds. Radiopharm., **1999**, 42, 589) with for example KCl, KI, KF or KCN gives a compound of formula (VIII)

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wherein R, R_1 and R_7 have the meanings given hereinabove and Nu represents a chloro, iodo, fluoro or cyano group. The reaction is preferably carried out in the presence of a weak base like NaHCO₃ or in the presence of a crown ether or a cryptand. (For more information on crown ethers and cryptands see: M. B. Smith and

J. March: Advanced organic chemistry, p. 105, 5th ed., (2001) John Wiley & Sons, New York, ISBN: 0-471-58589-0).

Compounds of general formula (VII) or (VIII) which have been obtained according to synthesis route D can be converted to compounds of general formula (I) analogously to the procedures described in synthesis route A, step 1 of route A, or step 2 of route A (see above).

Example 1

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10 Part A: To a 1M solution of sodium bis(trimethylsilyl) amide in THF (70 mL) is added dropwise a solution of 4-chloroaniline (8.86 gram, 69.5 mmol) in anhydrous THF in a nitrogen atmosphere. After the mixture is stirred for 20 minutes a solution of 2,4-dichlorobenzonitrile (12 gram, 70 mmol) in THF is added. The resulting mixture is stirred overnight, poured into ice-water (400 mL) and extracted with dichloromethane, dried over Na₂SO₄ and concentrated *in vacuo* to give a yellow oil (15.7 gram). Crystallisation from a dichloromethane/heptane mixture, and subsequent washing with methyl-t-butyl ether gives N-(4-chlorophenyl)-2,4-dichlorobenzenecarboxamidine (8.66 gram, 42 % yield) as a yellow solid. Meliting point (MP): 93-95 °C.

20 Analogously was prepared:

N-(4-bromophenyl)-2,4-dichlorobenzenecarboxamidine. MP: 117-119 °C.

Part B: A mixture of N-(4-chlorophenyl)-2,4-dichlorobenzenecarboxamidine (2.00 gram, 6.68 mmol), ethyl 3-bromo-2-oxopropanoate (2.65 gram, 13.6 mmol) and NaHCO₃ (1.12 gram, 13.3 mmol) in 2-propanol is stirred at reflux temperature for 20 hours. After cooling to room temperature the mixture is concentrated in vacuo and the residue suspended in dichloromethane, washed with water (3 x 50 mL) and brine (3 x 50 mL). The aqueous layers are extracted with dichloromethane. The combined organic layers are dried over Na₂SO₄ and concentrated in vacuo to afford crude brown product (2.0 gram). This product is further purified by column chromatography (silicagel, heptane/EtOAc = 90/10 (v/v)) to yield ethyl 1-(4-chlorophenyl)-2-(2,4dichlorophenyl)-1H-imidazole-4-carboxylate (0.759 gram, 29 % yield) as a yellow oil which slowly solidifies on standing. Melting point: 150-152 °C; MS: 395 (MH⁺). ¹H-NMR (400 MHz, CDCl₃): δ 7.91 (s, 1H), 7.49 (dd, J = 8 Hz, J = 2 Hz, 1H), 7.29-7.36 (m, 4H), 7.07 (dt, J = 8 Hz, J = 2 Hz, 2H), 4.44 (q, J = 7 Hz, 2H), 1.42 (t, J = 7 Hz, 3H).Part C: Ethyl 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate (0.810 gram, 2.06 mmol) and LiOH (0.173 g, 7.20 mmol) are dissolved in a H₂O/THF (20 mL/20 mL) mixture and stirred at 50 °C for 16 hours. The mixture is concentrated in vacuo to give 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylic acid. Thionyl chloride (60 mL) is added and the mixture is heated at reflux temperature for 1 hour and concentrated *in vacuo* to give crude 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carbonyl chloride.

Part D: Crude 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carbonyl chloride (919 mg, ~2.39 mmol), 1-aminopiperidine (0.469 g, 4.69 mmol) and triethylamine (0.363 g, 3.59 mmol) are dissolved in dichloromethane and stirred for one hour at room temperature. The mixture is washed with a saturated aqueous NaHCO₃ solution (3 x 20 mL), dried over Na₂SO₄ and concentrated *in vacuo* and further purified by column chromatography (ethyl acetate, silicagel) to give 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide

10 (356 mg, 26 % yield (based on ethyl 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate). Mass Spectrometry (MS): 449.

Analogously were prepared:

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- 15 2. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-(pyrrolidin-1-yl)-1H-imidazole-4-carboxamide; MS: 435.
 - 3. N-(t-Butoxy)-1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxamide; MS: 438.
 - 4. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-phenyl-1H-imidazole-4-carboxamide; MS: 442.
 - 5. 1-(4-Chlorophenyl)-N-cyclohexyl-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxamide; MS: 448.
 - 6. N-(Benzyl)-1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-N-methyl-1H-imidazole-4-carboxamide; MS: 470.
- 7. 1-[1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-4-(1H-imidazolyl)carbonyl] hexahydro-1H-azepine; MS: 448.
 - 8. 2-(4-Chlorophenyl)-1-(2,4-dichlorophenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide (prepared from 2,4-dichloroaniline and 4-chlorobenzo-nitrile); MS: 449.
- N-(t-Butoxy)-2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-1H-imidazole-4carboxamide (prepared from 2,4-dichloroaniline and 4-chlorobenzonitrile); MS: 438.

Example 10

Part A: Diisopropylamine (2.30 gram, 22.8 mmol) is added dropwise to anhydrous THF (100 mL) in a nitrogen atmosphere at 0 °C. n-BuLi is added dropwise (7.34 mL, 2.5 M solution in hexane, 18.4 mmol). The resulting solution is cooled to - 78 °C. A solution of ethyl 2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate (6.0 gram, 15.2 mmol) in anhydrous THF is added dropwise. The colour of the mixture changes from yellow to purple brown. The stirred mixture is warmed to - 40 °C and cooled to - 78 °C and allowed to stand for 30 minutes. Methyl iodide (6.44 gram, 45.4 mmol) is added dropwise and the resulting solution is stirred for 30

min at - 78 °C and then allowed to attain room temperature. The resulting solution is quenched with an aqueous NH₄Cl solution, diethyl ether is added and the organic layer is dried over MgSO₄, filtered and concentrated *in vacuo* to give an oil (6.4 gram). This oil is purified by column chromatography (toluene/EtOAc = 10/2 (v/v), silicagel) to give pure ethyl 2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxylate (5.3 gram, 85 % yield) as a yellow oil.

Part B: Ethyl 2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxylate (0.250 gram, 0.61 mmol) and LiOH (0.052 gram, 2.17 mmol) are dissolved in H_2O/THF (1:1 (v/v); 50 mL) and stirred at 50 °C for one hour. The mixture is concentrated to give crude 2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxylic acid. To this mixture is added $SOCl_2$ (50 mL) and the resulting mixture is heated at reflux temperature for 1 hour. The mixture is concentrated to give 2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carbonyl chloride.

Part C: 2-(4-Chlorophenyl)-1-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carbonyl chloride (1.5 gram, 3.75 mmol), 1-aminopiperidine (0.725 gram, 7.25 mmol) and triethylamine (0.549 gram, 5.44 mmol) are dissolved in dichloromethane and stirred for one hour at room temperature. The mixture is washed with a saturated aqueous NaHCO₃ solution, dried over Na₂SO₄ and concentrated *in vacuo* and further purified by column chromatography (heptane/ethyl acetate = 1/1 (v/v), silicagel) to give 2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide (0.220 gram, 13 % yield) as a white foam. MS: 463.

Analogously were prepared:

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- 11. N-(t-Butoxy)-2-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide: MS: 452.
- 12. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide: MS: 463; Melting point: 165-167 °C.
- 30 13. N-(t-Butoxy)-2-(2,4-dichlorophenyl)-1-(4-chlorophenyl)-5-methyl-1H-imidazole-4-carboxamide: MS: 452.
 - 14. N-(t-Butoxy)-1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-5-ethyl-1H-imidazole-4-carboxamide: Amorphous. MS: 468.
 - 15. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-ethyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide: MS: 477.
 - 16. 1-(4-Bromophenyl)-N-(t-butoxy)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide: Amorphous.
 - 17. 1-(4-Bromophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide: MP: > 204 °C. TLC (Silicagel, EtOAc) R_f = 0.3.
- 40 18. 1-(4-Bromophenyl)-N-(t-butoxy)-2-(2,4-dichlorophenyl)-5-ethyl-1H-imidazole-4-carboxamide: Amorphous. TLC (Silicagel, CH₂Cl₂/acetone = 9/1 (v/v)) R_f = 0.45.

- 19. 1-(4-Bromophenyl)-2-(2,4-dichlorophenyl)-5-ethyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide: MP: > 140 $^{\circ}$ C. TLC (Silicagel, EtOAc) R_f = 0.4.
- 20. 1-(4-Bromophenyl)-N-cyclohexyl-2-(2,4-dichlorophenyl)-5-ethyl-1H-imidazole-4-carboxamide: Melting point > 135-140 °C.
- 5 21. 1-(4-Bromophenyl)-2-(2,4-dichlorophenyl)-5-ethyl-N-(n-pentyl)-1H-imidazole-4-carboxamide: Syrup. TLC (Silicagel, CH₂Cl₂/acetone = 19/1 (v/v)) R_f = 0.4.

Example 22

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Part A: To a stirred solution of ethyl 1-(4-bromophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate (6.10 gram, 0.0139 mol) in THF (70 mL) is added LiOH (0.67 gram, 0.0278 mol) and water (70 mL). The resulting mixture is stirred for 16 hours at 50 °C to give a clear solution. After cooling to room temperature, HCl (1N solution, 28 mL) is added to give an oily precipitate which completely solidifies on continued stirring and addition of water (70 mL). The precipitate is collected by filtration, washed with water and dried *in vacuo* to give 1-(4-bromophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylic acid (4.92 gram, 86 % yield). Melting point: 138-142 °C.

Part B: To a stirred suspension of 1-(4-bromophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylic acid (1.23 gram, 2.99 mmol) in dry acetonitrile (40 mL) is successively added diisopropylethylamine (DIPEA) (1.15 mL, 6.6 mmol), O-benzotriazol-1-yl-N, N, N', N'-tetramethyluronium hexafluorophos-phate (HBTU) (1.36 gram, 3.6 mmol) and 1-aminopiperidine (0.39 mL, 3.6 mmol). After stirring for 16 hours, the resulting mixture is concentrated *in vacuo*. The residue is dissolved in ethylacetate and an aqueous NaHCO₃ solution is added. The ethylacetate layer is collected, washed with water and brine, dried over Na₂SO₄, filtered and concentrated *in vacuo* to give a crude solid. This solid is further purified by recrystallisation from acetonitrile to give 1-(4-bromophenyl)-2-(2,4-dichlorophenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide (830 mg, 56 % yield). Melting point: 219-221 °C.

30 Analogously were prepared:

- 23. N-(t-Butoxy)-1-(4-bromophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxamide. Amorphous. TLC (Silicagel, Et_2O) $R_f = 0.3$.
- 24. 1-(4-Bromophenyl)-2-(2,4-dichlorophenyl)-N-(pyrrolidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 238-240 °C.
- 25. N-(Azepan-1-yl)-1-(4-bromophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 201-204 °C.
- 26. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-(hexahydrocyclopenta[c]pyrrol 2(1H)-yl)-1H-imidazole-4-carboxamide. MS: 475.
- 40 27. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-(4-fluorobenzyl)-1H-imidazole-4-carboxamide. MS: 474.

- 28. 1-(4-Chlorophenyl)-2-(2-methoxy-4-chlorophenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 220 °C.
- 29. 1-(4-Chlorophenyl)-N-cyclohexyl-2-(2-methoxy-4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 177-179 °C.
- 5 30. 1-(4-Chlorophenyl)-2-(2-fluoro-4-chlorophenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 217-218 °C. .
 - 31. 2-(2,4-Dichlorophenyl)-1-(4-fluorophenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 175-176 °C.
 - 32. N-Cyclohexyl-2-(2,4-dichlorophenyl)-1-(4-fluorophenyl)-1H-imidazole-4-carboxamide. Melting point: 184-185 °C.
 - 33. N-Cyclohexyl-2-(2-fluoro-4-chlorophenyl)-1-(4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 157-159 °C.
 - 34. 1-(4-Chlorophenyl)-2-(2-methoxy-4-chlorophenyl)-N-(n-pentyl)-1H-imidazole-4-carboxamide. Melting point: 115 °C.
- 15 35. 2-(2,4-Dichlorophenyl)-1-(4-methoxyphenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 178-179 °C.
 - 36. N-Cyclohexyl-2-(2,4-dichlorophenyl)-1-(4-methoxyphenyl)-1H-imidazole-4-carboxamide. Melting point: 175-176 °C.
 - 37. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N,N-diethyl-1H-imidazole-4-carboxamide. Melting point: 177-179 °C.
 - 38. 1-(4-Chlorophenyl)-N-cyclohexyl-2-(2-trifluoromethyl-4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 172 °C.
 - 39. 1-(4-Chlorophenyl)-N-(piperidin-1-yl)-2-(2-trifluoromethyl-4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 219 °C.
- 40. N-(1-Adamantyl)-1-(4-chlorophenyl)-2-(2-trifluoromethyl-4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 288 °C.
 - 41. 1-(4-Chlorophenyl)-N-(2,2,2-trifluoroethyl)-2-(2-trifluoromethyl-4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 149 °C.
 - 42. 2-(2,4-Dichlorophenyl)-1-(pyridin-3-yl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 165-170 °C.
 - 43. N-Cyclohexyl-2-(2,4-dichlorophenyl)-1-(pyridin-3-yl)-1H-imidazole-4-carboxamide.Melting point: 195 °C.
 - 44. 2-(2,4-Dichlorophenyl)-1-(pyridin-3-yl)-N-(n-pentyl)-1H-imidazole-4-carboxamide. Melting point: 117 °C.

Example 45

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Part A: 2,4-Dichlorobenzoyl chloride (40.0 g, 0.19 mol) is dissolved in tetrahydrofuran (1 L). To the resulting stirred solution is successively added disopropylethylamine (DIPEA) (73.4 mL, 2.2 molar equivalent) and 4-(trifluoromethyl)phenylamine (30.7 g, 0.19 mol). After one hour the mixture is concentrated *in vacuo* to give an oil. This oil is crystallised from ethanol to give pure 2,4-dichloro-N-(4-(trifluoromethyl)phenyl)benzamide (53.2 g, 83 % yield). ¹H-NMR

(200 MHz, DMSO-d₆): δ 10.90 (br s, 1H), 7.91 (br d, J = 8 Hz, 2H), 7.63-7.77 (m, 4H), 7.57 (dt, J = 8 Hz, J= 2 Hz, 1H).

Part B: 2,4-Dichloro-N-(4-(trifluoromethyl)phenyl)benzamide (19.0 g, 0.057 mol) is dissolved in benzene (150 mL) and PCl₅ (13.0 g, 1.1 molar equivalent) is added. The resulting mixture is heated at reflux temperature for two hours, allowed to attain room temperature and concentrated *in vacuo* to give a residue. The residue is dissolved in anhydrous THF, cooled to 0 °C and transferred into an autoclave. Excess NH₃ is quickly added from a lecture bottle and the mixture is stirred at room temperature for 50 hours. A mixture of ethylacetate and aqueous NaHCO₃ is added. The ethylacetate layer is collected, dried over Na₂SO₄, filtered and concentrated *in vacuo*. The resulting oil is purified by column chromatography (diethyl ether/petroleum ether = 1/1 (v/v), silicagel) to give pure 2,4-dichloro-N-(4-(trifluoromethyl)phenyl)benzene-carboxamidine (16.9 g, 89 % yield). Melting point: 108-109 °C.

Part C: 2,4-Dichloro-N-(4-(trifluoromethyl)phenyl)benzenecarboxamidine (15.0 g, 0.0450 mol) is dissolved in 2-propanol and ethyl 3-bromo-2-oxobutanoate (20.8 g, 2 molar equivalent) and NaHCO₃ are successively added. The resulting mixture is heated at reflux temperature for 40 hours and allowed to attain room temperature. The 2-propanol is removed *in vacuo*, ethyl acetate is added to the residue and the resulting organic layer is washed with NaHCO₃ (5 % aqueous solution). The ethylacetate layer is collected, dried over Na₂SO₄, filtered and concentrated *in vacuo*. The resulting oil is purified by column chromatography (diethyl ether/petroleum ether = 1/3 (v/v), silicagel) and further purified by crystallisation from cyclohexane to give ethyl 2-(2,4-dichlorophenyl)-5-methyl-1-(4-(trifluoromethyl)phenyl)-1H-imidazole-4-carboxylate (10.45 g, 52 % yield) as a yellow solid. Melting point: 160-162 °C.

Part D: The formed ethyl 2-(2,4-dichlorophenyl)-5-methyl-1-(4-(trifluoromethyl) phenyl)-1H-imidazole-4-carboxylate is converted to 2-(2,4-dichlorophenyl)-5-methyl-1-(4-(trifluoromethyl)phenyl)-1H-imidazole-4-carboxylic acid (melting point: 224-226 °C), which carboxylic acid is converted to 2-(2,4-dichlorophenyl)-5-methyl-N-(piperidin-1-yl)-1-(4-(trifluoromethyl)phenyl)-1H-imidazole-4-carboxamide (melting point: 173-174 °C) according to the procedure described in example 22 above. Analogously were prepared

- 46. 2-(2,4-dichlorophenyl)-N-(piperidin-1-yl)-1-(4-(trifluoromethyl)phenyl)-1H-imidazole-4-carboxamide. Melting point: >200 °C (decomposition).
- 47. N-Cyclohexyl-2-(2,4-dichlorophenyl)-5-methyl-1-(4-(trifluoromethyl) phenyl)-1H-imidazole-4-carboxamide. Melting point: 178-179 °C.
 - 48. N-Cyclohexyl-2-(2,4-dichlorophenyl)-1-(4-(trifluoromethyl)phenyl)-1H-imidazole-4-carboxamide. Melting point: 199-200 °C.

40 <u>Example 49</u>

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Part A: N-(4-methoxyphenyl)-2,4-dichlorobenzenecarboxamidine (15.0 gram, 50.8 mmol) is dissolved in 2-propanol and ethyl 3-bromo-2-oxobutanoate (23.5 g, 2 molar

equivalents) and NaHCO₃ (8.5 gram, 2 molar equivalents) are successively added. The resulting mixture is heated at reflux temperature for 40 hours and allowed to attain room temperature. The 2-propanol is removed *in vacuo*, ethyl acetate is added to the residue and the resulting organic layer is washed with NaHCO₃ (5 % aqueous solution). The ethylacetate layer is collected, dried over Na₂SO₄, filtered and concentrated *in vacuo*. The resulting oil is purified by column chromatography (diethyl ether/petroleum ether = 1/3 (v/v), silicagel) to give ethyl 2-(2,4-dichlorophenyl)-5-methyl-1-(4-methoxy-phenyl)-1H-imidazole-4-carboxylate (8.61 g, 42 % yield) as a solid. 1 H-NMR (200 MHz, CDCl₃): δ 7.33 (d, J = 8 Hz, 1H), 7.27 (d, J = 2 Hz, 1H), 7.18 (dd, J = 8 Hz, J = 2 Hz, 1H), 7.03 (dt, J = 8 Hz, J = 2 Hz, 2H), 6.85 (dt, J = 8 Hz, J = 2 Hz, 2H), 4.42 (q, J = 7 Hz, 2H), 3.80 (s, 3H), 2.43 (s, 3H), 1.43 (t, J = 7 Hz, 3H).

Part B: To a stirred solution of ethyl 2-(2,4-dichlorophenyl)-5-methyl-1-(4-methoxyphenyl)-1H-imidazole-4-carboxylate (8.00 gram, 0.0198 mol) in THF (80 mL) is added LiOH (0.59 gram, 2 molar equivalents) and water (80 mL). The resulting mixture is stirred for 16 hours at 80 °C. After cooling to room temperature, HCl (2N solution, 12.3 mL) is added to give an oily precipitate. After addition of water and extraction with ethylacetate, the ethylacetate layer is collected, dried over Na₂SO₄, filtered and concentrated *in vacuo*. The residue is crystallised from diisopropyl ether and dried to give 2-(2,4-dichlorophenyl)-5-methyl-1-(4-methoxyphenyl)-1H-imidazole-4-carboxylic acid (4.04 gram, 87 % yield) as a pale grey solid. Melting point: 189-191 °C.

Part C: To 2-(2,4-dichlorophenyl)-5-methyl-1-(4-methoxyphenyl)-1H-imidazole-4-carboxylic acid (1.00 gram, 2.65 mmol) in dry acetonitrile (25 mL) is successively added diisopropylethylamine (DIPEA) (1.02 mL, 2.2 molar equivalents), O-benzotriazol-1-yl-N, N, N', N'-tetramethyluronium hexafluoro-phosphate (HBTU) (1.21 gram, 1.2 molar equivalents) and the resulting solution is stirred for 15 minutes. Cyclohexylamine (0.36 mL, 1.2 molar equivalents) is added. After stirring for 50 hours, the resulting mixture is concentrated *in vacuo*. The residue is dissolved in dichloromethane and an aqueous NaHCO₃ solution is added. The dichloromethane layer is collected, dried over Na₂SO₄, filtered and concentrated *in vacuo*. The residue is further purified by column chromatography (gradient: dichloromethane => dichloromethane/methanol = 99/1 (v/v), silicagel) to give N-(1-cyclohexyl)-2-(2,4-dichlorophenyl)-5-methyl-1-(4-methoxyphenyl)-1H-imidazole-4-carboxamide (1.03 gram, 85 % yield). Melting point: 160-161 °C.

Analogously were prepared:

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- 50. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N,N,5-trimethyl-1H-imidazole-4-carboxamide. Melting point: 101-104 °C.
- 51. 1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. MS: 464 (MH⁺).

- 52. 1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-methyl-N-(4-morpholinyl)-1H-imidazole-4-carboxamide. MS: 466 (MH⁺).
- 53. N-(1-Azepanyl)-1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. MS: 478 (MH⁺).
- 5 54. 1-(4-Chloropyridin-2-yl)-N-cyclohexyl-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. MS: 463.
 - 55. 1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-methyl-N-(n-pentyl)-1H-imidazole-4-carboxamide. MS: 451.
 - 56. 1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-N-(4-fluorobenzyl)-5-methyl-1H-imidazole-4-carboxamide. MS: 489. Melting point: 123-126 °C.
 - 57. 1-(4-Chlorophenyl)-N-cyclohexyl-5-methyl-2-(2-trifluoromethyl-4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 212 °C.
 - 58. 1-(4-Chlorophenyl)-5-methyl-N-(piperidin-1-yl)-2-(2-trifluoromethyl-4-chlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 165 °C.

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- 15 59. 1-(4-Chlorophenyl)-2-(2-methoxy-4-chlorophenyl)-5-methyl-N-(n-pentyl)-1H-imidazole-4-carboxamide. Melting point: 131 °C.
 - 60. 1-(4-Chlorophenyl)-2-(2-methoxy-4-chlorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: > 256 °C.
 - 61. N-Cyclohexyl-1-(4-chlorophenyl)-2-(2-methoxy-4-chlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 201 °C.
 - 62. 2-(2,4-Dichlorophenyl)-1-(4-fluorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 223-224 °C.
 - 63. 2-(2,4-Dichlorophenyl)-5-methyl-1-(4-methoxyphenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: > 90 °C (decomposition).
- 25 64. N-Cyclohexyl-1-(4-fluorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 229-230 °C.
 - 65. 1-(4-Chlorophenyl)-5-methyl-N-(n-pentyl)-2-(2-trifluoromethyl-4-chlorophenyl)-1H-imidazole-4-carboxamide. Amorphous.
 - 66. 1-(4-Chlorophenyl)-2-(2-fluoro-4-chlorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 195 °C.
 - 67. 1-(4-Chlorophenyl)-2-(2-fluoro-4-chlorophenyl)-5-methyl-N-(n-pentyl)-1H-imidazole-4-carboxamide. Melting point: 115 °C.
 - 68. 1-(4-Chlorophenyl)-N-(cyclohexyl)-2-(2-fluoro-4-chlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 188 °C.
- 35 69. 1-(4-Chlorophenyl)-N-(cyclohexyl)-2-(1,5-dimethyl-1H-pyrrol-2-yl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 188-189 °C.
 - 70. 1-(4-Chlorophenyl)-2-(1,5-dimethyl-1H-pyrrol-2-yl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 208-210 °C.
 - 71. 2-(2-Chlorophenyl)-1-(3-fluorophenyl)-5-methyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 236-238 °C.
 - 72. 2-(2-Chlorophenyl)-1-(3-fluorophenyl)-5-methyl-N-(n-pentyl)-1H-imidazole-4-carboxamide. Melting point: 97-102 °C.

- 73. 2-(2-Chlorophenyl)-N-cyclohexyl-1-(3-fluorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 180-182.5 °C.
- 74. 2-(2-Chlorophenyl)-1-(3-fluorophenyl)-N-(2-(4-fluorophenyl)ethyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 123.5-126 °C.
- 5 75. 1-(4-Chloropyridin-2-yl)-2-(2;4-dichlorophenyl)-5-ethyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 146 °C.
 - 76. 1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-ethyl-N-(4-morpholinyl)-1H-imidazole-4-carboxamide. Melting point: 223 °C.
 - 77. N-(1-Azepanyl)-1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-ethyl-1H-imidazole-4-carboxamide. Melting point: 177 °C.
 - 78. 1-(4-Chloropyridin-2-yl)-N-cyclohexyl-2-(2,4-dichlorophenyl)-5-ethyl-1H-imidazole-4-carboxamide. Melting point: 149 °C.

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- 79. 1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-ethyl-N-(n-pentyl)-1H-imidazole-4-carboxamide. Melting point: Oil.
- 15 80. 1-(4-Chloropyridin-2-yl)-2-(2,4-dichlorophenyl)-5-ethyl-N-(4-fluorophenylmethyl)-1H-imidazole-4-carboxamide. MP: amorphous.
 - 81. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-(hexahydrocyclopenta-[c]pyrrol-2(1H)-yl)-5-methyl-1H-imidazole-4-carboxamide. MP: 143-146 °C.
 - 82. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-phenyl-1H-imidazole-4-carboxamide. Melting point: 91-95 °C.
 - 83. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(tetrahydro-2H-pyran-2-yloxy)-1H-imidazole-4-carboxamide. Melting point: 128-133 °C.
 - 84. N-(Exo-bicyclo[2.2.1]hept-2-yl)-1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 194-195 °C.
- 25 85. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-(2-fluoroethyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 128-133 °C.
 - 86. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-(trans-4-hydroxycyclohexyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 160 °C (dec.).
 - 87. 1-{[1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazol-4-yl]carbonyl}-4-hydroxypiperidine. Melting point: Amorphous.
 - 88. 1-{[1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazol-4-yl]carbonyl}-1,2,3,4-tetrahydroisoquinoline. Melting point: 143-146 °C.
 - 89. N-(Endo-bicyclo[2.2.1]hept-2-yl)-1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 194-195 °C.
- 35 90. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-N-(4-fluorobenzyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 165-166 °C.
 - 91. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(n-pentyl)-1H-imidazole-4-carboxamide. Oil.
 - 92. N-(Azepan-1-yl)-1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 147-149 °C.
 - 93. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(pyrrolidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 205-206 °C.

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- 94. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(morpholin-4-yl)-1H-imidazole-4-carboxamide. Melting point: 225 °C (dec.).
- 95. 2-(2,5-Dichlorophenyl)-5-methyl-1-phenyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 227 °C.
- 5 96. N-Cyclohexyl-2-(2,5-dichlorophenyl)-5-methyl-1-phenyl-1H-imidazole-4-carboxamide. Melting point: 236 °C.
 - 97. N-Cyclohexyl-2-(2,4-dichlorophenyl)-1-(2,5-difluorophenyl)-5-ethyl-1H-imidazole-4-carboxamide. Melting point: 144-146 °C.
 - 98. N-Cyclohexyl-2-(2,4-dichlorophenyl)-1-(2,5-difluorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 206-208 °C.
 - 99. N-Cyclohexyl-2-(1,5-dimethyl-1H-pyrrol-2-yl)-5-ethyl-1-phenyl-1H-imidazole-4-carboxamide. Melting point: 195-196 °C.
 - 100. N-Cyclohexyl-2-(2,5-dichlorophenyl)-5-ethyl-1-phenyl-1H-imidazole-4-carboxamide. Melting point: 198-199 °C.
- 15 101. 2-(2,5-Dichlorophenyl)-5-ethyl-1-phenyl-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 207-208 °C.
 - 102. 1-(4-Chlorophenyl)-5-methyl-2-(3-methylpyridin-2-yl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 211-213 °C.
 - 103. 1-(4-Chlorophenyl)-N-cyclohexyl-5-methyl-2-(3-methylpyridin-2-yl)-1H-imidazole-4-carboxamide. Melting point: 188-190 °C.
 - 104.1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(3-(trifluoromethyl)phenyl)-1H-imidazole-4-carboxamide. Melting point: 177 °C.
 - 105.1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(3-(trifluoromethyl)benzyl)-1H-imidazole-4-carboxamide. Melting point: 138-140 °C.
- 25 106.1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-N-(4- (trifluoromethyl)benzyl)-1H-imidazole-4-carboxamide. Melting point: 232 °C.
 - 107.1-(4-Chlorophenyl)-N-cyclopentyl-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxamide. Melting point: 172 °C.
- 108.1-(4-Chlorophenyl)-N-cycloheptyl-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-30 4-carboxamide. Melting point: 154-156 °C.

Example 109

- **Part A**: Ethyl 1-(4-bromophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate is converted to ethyl 1-(4-bromophenyl)-5-chloro-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate analogously to a published procedure (N. Kudo et al., *Chem. Pharm. Bull.* **1999**, *47*, 857-868) using excess of SO₂Cl₂ in dichloroethane at reflux temperature for 50 hours.
- Part B: Ethyl 1-(4-bromophenyl)-5-chloro-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate is converted to 1-(4-bromophenyl)-5-chloro-2-(2,4-dichloro-phenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide (melting point: > 150 °C; R_f (Silicagel, EtOAc) ~ 0.35) analogously to the procedure described in example 22 above. 1 H-NMR (400 MHz, CDCl₃): δ 7.85 (br s, 1H), 7.52 (dt, J = 8 Hz, J = 2 Hz, 2H), 7.26-7.36

(m, 3H), 7.01 (dt, J = 8 Hz, J = 2 Hz, 2H), 2.85-2.92 (m, 4H), 1.72-1.80 (m, 4H), 1.40 - 1.44 (m, 2H).

Example 110

Part A: To a stirred solution of 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylic acid (18.38 gram, 50 mmol) in toluene (200 mL) in a nitrogen atmosphere is added N,N-dimethylformamide di-tert-butyl acetal (50 mL) and the resulting mixture is heated at 80 °C for 4 hours. After cooling to room temperature the reaction mixture is concentrated and diethyl ether is added. The resulting solution is twice washed with water, dried over MgSO₄, filtered and concentrated *in vacuo*. The residue is crystallised from diisopropyl ether to give pure tert-butyl 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylate (10.35 gram, 49 % yield). Melting point: 179-181 °C.

Part B:

Lithium diisopropyl amide (LDA) (5.25 mL of a 2 M solution in THF, 0.0105 mol) is 15 added dropwise to a cooled solution (-70 °C) of tert-butyl 1-(4-chlorophenyl)-2-(2,4dichlorophenyl)-1H-imidazole-4-carboxylate (4.24 gram, 0.010 mol) in anhydrous THF (80 mL) in a nitrogen atmosphere and the resulting mixture is stirred for one hour. A solution of p-toluenesulfonyl cyanide (1.88 gram, 0.011 mol) in anhydrous 20 THF (20 mL) is added dropwise and the resulting red solution is stirred for one hour at - 70 °C and then allowed to attain room temperature. Diethyl ether is added and the resulting solution is quenched with water and filtered over hyflo. The organic layer is collected and washed with water, dried over MgSO4, filtered and concentrated in vacuo to give an oil. This oil is purified by column chromatography (dichloromethane, 25 silicagel) to give 3.4 gram of tert-butyl 1-(4-chlorophenyl)-5-cyano-2-(2,4dichlorophenyl)-1H-imidazole-4-carboxylate. Recrystallisation from diisopropyl ether gave crystalline tert-butyl 1-(4-chlorophenyl)-5-cyano-2-(2,4-dichlorophenyl)-1Himidazole-4-carboxylate (2.57 gram, 57 % yield). Melting point: 210-212 °C.

30 Analogously was prepared:

Tert-butyl 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-5-methyl-1H-imidazole-4-carboxylate. ¹H-NMR (400 MHz, CDCl₃): δ 7.38 (d, J = 8 Hz, 1H), 7.34 (dt, J = 8 Hz, J = 2 Hz, 2H), 7.27 (d, J = 2 Hz, 1H), 7.22 (dd, J = 8 Hz, J = 2 Hz, 1H), 7.03 (dt, J = 8 Hz, J = 2 Hz, 2H), 2.40 (s, 3H), 1.63 (s, 9H).

Part C:

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To a solution of tert-butyl 1-(4-chlorophenyl)-2-(2,4-dichlorophenyl)-5-cyano-1H-imidazole-4-carboxylate (2.57 gram, 5.73 mmol) in dichloromethane (40 mL) is added trifluoroacetic acid and the resulting solution is stirred at room temperature for 20 hours and concentrated *in vacuo*. The residue is crystallised from diisopropyl ether to give pure 1-(4-chlorophenyl)-5-cyano-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylic acid (1.95 gram, 87 % yield). Melting point: 200-202 °C (dec.).

Part D:

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1-(4-Chlorophenyl)-5-cyano-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxylic acid is converted to 1-(4-chlorophenyl)-5-cyano-2-(2,4-dichlorophenyl)-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide in 60 % yield, analogously to the procedure described in example 22, part B herein above. Melting point: 231-233.5 °C.

Analogously were prepared:

- 111. 1-(4-Chlorophenyl)-2-(2,4-dichlorophenyl)-5-iodo-N-(piperidin-1-yl)-1H-imidazole-4-carboxamide. Melting point: 196-201 °C.
- 112. 1-(4-Chlorophenyl)-N-cyclohexyl-2-(2,4-dichlorophenyl)-5-iodo-1H-imidazole-4-carboxamide. Melting point: 226-230 °C.
- 113. 1-(4-Chlorophenyl)-5-cyano-N-cyclohexyl-2-(2,4-dichlorophenyl)-1H-imidazole-4-carboxamide. Melting point: 157-158 °C.

Claims

1. A compound of formula (I)

wherein

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- R represents phenyl, thienyl, 2-pyridinyl, 3-pyridinyl, 4-pyridinyl, pyrimidinyl, pyrazinyl, pyridazinyl or triazinyl, which groups may be substituted with 1, 2, 3 or 4 substituents Y, which can be the same or different, from the group C₁₋₃-alkyl or alkoxy, hydroxy, halogen, trifluoromethyl, trifluoromethylthio, trifluoromethoxy, nitro, amino, mono- or dialkyl (C₁₋₂)-amino, mono- or dialkyl (C₁₋₂)-amido, (C₁₋₃)-alkoxycarbonyl, carboxyl, cyano, carbamoyl and acetyl, or R represents naphtyl, with the proviso that when R is 4-pyridinyl, R₄ represents a halogen atom or a cyano, carbamoyl, formyl, acetyl, trifluoroacetyl, fluoroacetyl, propionyl, sulfamoyl, methanesulfonyl, methylsulfanyl or branched or unbranched C₁₋₄ alkyl group, which C₁₋₄ alkyl group may be substituted with 1-3 fluoro atoms or with a bromo, chloro, iodo, cyano or hydroxy group,
- R₁ represents phenyl or pyridinyl, which groups may be substituted with 1-4 substituents Y, which can be the same or different, wherein Y has the above mentioned meaning, or R₁ represents pyrimidinyl, pyrazinyl, pyridazinyl or triazinyl, which groups may be substituted with 1-2 substituents Y, which can be the same or different or R₁ represents a five-membered aromatic heterocyclic ring having one or two heteroatoms from the group (N, O, S), which heteroatoms can be the same or different, which five-membered aromatic heterocyclic ring may be substituted with 1-2 substituents Y, which can be the same or different or R₁ represents naphtyl,
 - R₂ represents H, branched or unbranched C₁₋₈ alkyl, C₃₋₈ cycloalkyl, C₃₋₈ alkenyl,
 C₅₋₈ cycloalkenyl which groups may contain a sulfur, oxygen or nitrogen atom,
- R₃ represents branched or unbranched C₂₋₈ alkyl, C₁₋₈ alkoxy, C₅₋₈ cycloalkyloxy, C₃₋₈ cycloalkyl, C₅₋₁₀ bicycloalkyl, C₆₋₁₀ tricycloalkyl, C₃₋₈ alkenyl, C₅₋₈ cycloalkenyl, which groups may optionally contain one or more heteroatoms from the group (O, N, S) and which groups may be substituted with a hydroxy group or 1-2 C₁₋₃ alkyl groups or 1-3 fluoro atoms, or R₃ represents a benzyl or phenethyl group which aromatic rings may be substituted with 1-5 substituents Z, which can be the same or different, from the group C₁₋₃-alkyl or alkoxy, hydroxy, halogen, trifluoromethyl, trifluoromethylthio, trifluoromethoxy, nitro, amino, mono- or dialkyl (C₁₋₂)-amino, mono- or dialkyl (C₁₋₂)-amido, (C₁₋₃)-alkylsulfonyl, dimethyl-sulfamido, C₁₋₃-

alkoxycarbonyl, carboxyl, trifluoromethylsulfonyl, cyano, carbamoyl, sulfamoyl and acetyl, or R_3 represents a phenyl or pyridinyl group, which groups are substituted with 1-4 substituents Z, wherein Z has the meaning as indicated above,

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- or R₃ represents a pyridinyl group, or R₃ represents a phenyl group, with the proviso that R₄ represents a halogen atom or a cyano, carbamoyl, formyl, acetyl, trifluoroacetyl, fluoroacetyl, propionyl, sulfamoyl methanesulfonyl, methylsulfanyl or C₁₋₄ alkyl group, which C₁₋₄ alkyl group may be substituted with 1-3 fluoro atoms or with a bromo, chloro, iodo, cyano or hydroxy group,
- or R₃ represents a group NR₅R₆ with the proviso that R₂ represents a hydrogen atom or a methyl group, wherein
- R₅ and R₆ are the same or different and represent branched or unbranched C₁₋₄ alkyl, or R₅ and R₆ together with the nitrogen atom to which they are bonded form a saturated or unsaturated, monocyclic or bicyclic heterocyclic group having 4 to 10 ring atoms which heterocyclic group contains one or two heteroatoms from the group (N, O, S), which heteroatoms can be the same or different, which heterocyclic group may be substituted with a C₁₋₃ alkyl group or a hydroxy group, or R₂ and R₃ together with the nitrogen atom to which they are bonded form a saturated or unsaturated heterocyclic group having 4 to 10 ring atoms which heterocyclic group contains one or two heteroatoms from the group (N, O, S), which heteroatoms can be the same or different, which heterocyclic group may be substituted with a C₁₋₃ alkyl group or a hydroxy group,
 - R₄ represents a hydrogen or halogen atom or a cyano, carbamoyl, formyl, acetyl, trifluoroacetyl, fluoroacetyl, propionyl, sulfamoyl, methanesulfonyl, methylsulfanyl or branched or unbranched C₁₋₄ alkyl group, which C₁₋₄ alkyl group may be substituted with 1-3 fluoro atoms or with a bromo, chloro, iodo, cyano or a hydroxy group,

and prodrugs, stereoisomers and salts thereof.

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- 2. Pharmaceutical compositions containing a pharmacologically active amount of at least one compound as claimed in 1 as an active component.
- Method of preparing pharmaceutical compositions as claimed in claim 2
 characterised in that a compound as claimed in claim 1 is brought in a form suitable for administration.
 - 4. Process for the preparation of compounds having formula (I), characterised in that a compound is prepared wherein R, R₁-R₃ have the meanings given in claim 1 and R₄ represents a hydrogen or halogen atom or a cyano, carbamoyl, formyl, acetyl, trifluoroacetyl, propionyl, sulfamoyl, methanesulfonyl, methylsulfanyl or C₁₄ alkyl group, which C₁₄ alkyl group may be substituted with 1-3 fluoro atoms,

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by reacting a compound having formula (II), (III) or (IV) with a compound of formula R_2R_3NH .

5. Process for the preparation of compounds having formula (II)

$$R_1$$
 N
 R_4
 R_4
 R
 R
 R
 R
 R
 R
 R

wherein R_4 represents a C_{1-4} alkyl group, which C_{1-4} alkyl group may be substituted with 1-3 fluoro substituents, or wherein R_4 represents a halogen atom or a cyano, formyl, acetyl, trifluoroacetyl, fluoroacetyl, methylsulfanyl or propionyl group, characterized in that a compound is prepared wherein R and R_1 have the meanings given in claim 1 and R_7 represents a branched or unbranched alkyl group (C_{1-4}) or benzyl group, by reacting a compound having formula (II) wherein R_4 is a hydrogen atom with a compound having formula R_4 -X, wherein X represents a leaving group and R_4 represents a C_{1-4} alkyl group, which C_{1-4} alkyl group may be substituted with 1-3 fluoro substituents, or wherein R_4 represents a halogen atom or a cyano, formyl, acetyl, trifluoroacetyl, fluoroacetyl, methylsulfanyl or propionyl group, in the presence of a strong non-nucleophilic base.

6. Process for the preparation of a compound having formula (II)

$$R_1$$
 N
 R_4
 R
 R
 R
 R

wherein R₄ represents a branched or unbranched C₁₋₄ alkyl group, which C₁₋₄ alkyl group may be substituted with 1-3 fluoro substituents, characterized in that a compound is prepared wherein R, R₁ have the meanings given in claim 1 and R₇ represents a branched or unbranched alkyl group (C₁₋₄) or benzyl group, by reacting a compound having formula (V) or its tautomer

wherein R and R_1 have the meanings given in claim 1, with a compound having formula (VI)

$$R_4$$
 O
 OR_7
 OR_7
 OR_7

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wherein R_4 represents a branched or unbranched C_{1-4} alkyl group, which C_{1-4} alkyl group may be substituted with 1-3 fluoro atoms and R_8 represents a so-called leaving group and R_7 represents a branched or unbranched alkyl group (C_{1-4}) or benzyl group.

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7. Compounds of formula (IX)

$$R_1 \xrightarrow[R]{N} R_4$$
(IX)

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wherein R and R_4 have the meanings given in claim 1 and wherein R_1 represents a phenyl or pyridinyl group, which groups are substituted with 1-4 substituents Y, which can be the same or different, or R_1 represents a pyrimidinyl, pyrazinyl, pyridazinyl or triazinyl group, which groups are substituted with 1-2 substituents Y, which can be the same or different or R_1 represents a five-membered aromatic heterocyclic moiety having one or two heteroatoms from the group (N, O, S), which heteroatoms can be the same or different, which five-membered aromatic heterocyclic moiety may be substituted with 1-2 substituents Y, which can be the same or different or R_1 represents naphtyl and R_9 represents a hydroxy group, a branched or unbranched alkoxy (C_{1-4}) group, a benzyloxy group or a chloro substituent.

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8. Compounds of formula (X) and tautomers thereof

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wherein R represents a 4-chlorophenyl group, a 4-bromophenyl group or a 4-(trifluoromethyl)phenyl group.

Use of a compound as claimed in claim 1 for the preparation of a pharmaceutical composition for the treatment of disorders involving cannabinoid neurotransmission.

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10. Use as claimed in claim 9 characterised in that said disorders are psychiatric disorders such as psychosis, anxiety, depression, attention deficits, memory disorders, cognitive disorders, appetite disorders, obesity, addiction, appetence, drug dependence and neurological disorders such as neurodegenerative disorders, dementia, dystonia, muscle spasticity, tremor, epilepsy, multiple sclerosis, traumatic brain injury, stroke, Parkinson's disease, Alzheimer's disease, epilepsy, Huntington's disease, Tourette's syndrome, cerebral ischaemia, cerebral apoplexy, craniocerebral trauma, stroke, spinal cord injury, neuroinflammatory disorders, plaque sclerosis, viral encephalitis, demyelinisation related disorders, as well as for the treatment of pain disorders, including neuropathic pain disorders, and other diseases involving cannabinoid neurotransmission, including the treatment of septic shock, glaucoma, cancer, diabetes, emesis, nausea, asthma, respiratory diseases, gastrointestinal disorders, gastric ulcers, diarrhoea and cardiovascular disorders.

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- (71) Applicant (for all designated States except US): SOLVAY PHARMACEUTICALS B.V. [NL/NL]; C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): KRUSE, Cornelis, G. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL). LANGE, Josephus, H.M. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL). HERREMANS, Arnoldus, H.J. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL). VAN STUIVENBERG, Herman, H. [NL/NL]; Solvay Pharmaceuticals B.V., c/o C.J. van Houtenlaan 36, NL-1381 CP Weesp (NL).

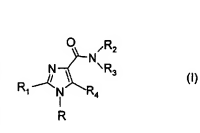
- (74) Agent: MUIS, Maarten; OCTROOIBUREAU ZOAN B.V., P.O. Box 140, NL-1380 AC Weesp (NL).
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(54) Title: 1H-IMIDAZOLE DERIVATIVES HAVING CB1 AGONISTIC, CB1 PARTIAL AGONISTIC OR CB1- ANTAGONIS-WO 03/027076 A TIC ACTIVITY



(57) Abstract: AbstractThe present invention relates to a group of novel 1H-imidazole derivatives, to methods for the preparation of these compounds, and to pharmaceutical compositions containing one or more of these compounds as an active component. These 1H-imidazole derivatives are potent cannabinoid-CB1 receptor agonists, partial agonists or antagonists, useful for the treatment of psychiatric and neurological disorders, as well as and other diseases involving cannabinoid neurotransmission. The compounds have the general formula (I) wherein R and R1-R4 have the meanings given in the specification.

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A. CLASSIFICATION OF SUBJECT MATTER
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According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

 $\begin{array}{ccc} {\rm Minimum\ cocumentation\ searched\ (classification\ system\ followed\ by\ classification\ symbols)} \\ {\rm IPC\ 7\ C070\ C07C\ A61K} \end{array}$

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

BEILSTEIN Data, WPI Data, PAJ, CHEM ABS Data, EPO-Internal

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χ Funt	her documents are listed in the continuation of box C.	Patent family members are liste	ed in annex.	
"A" docume consid "E" earlier of filing d "L" docume which citation "O" docume other r	negories of cited documents: ant defining the general state of the art which is not lered to be of particular relevance document but published on or after the international state and which may throw doubts on priority daim(s) or is cited to establish the publication date of another nor other special reason (as specified) and referring to an oral disclosure, use, exhibition or means and published prior to the International filing date but not the priority date claimed	"T" later document published after the in or priority date and not in conflict will cited to understand the principle or invention "X" document of particular relevance; the cannot be considered novel or canninvolve an inventive step when the "Y" document of particular relevance; the cannot be considered to involve an document is combined with one or ments, such combined with one or ments, such combination being obvin the act. "å" document member of the same pate	in the application but theory underlying the claimed invention to be considered to document is taken alone claimed invention step when the more other such docu-ious to a person skilled	
Date of the	actual completion of the international search	Date of mailing of the international	search report	
4	December 2002	11/12/2002		
Name and r	mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax (+31-70) 340-3018	Authorized officer Scruton-Evans, I		

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Category °	Citation of document, with Indication, where appropriate, of the relevant passages	Relevant to claim No.	
Y	WO 00 46209 A (SANOFI SYNTHELABO ;BARTH FRANCIS (FR); CAMUS PHILIPPE (FR); MARTIN) 10 August 2000 (2000-08-10) the whole document	1-10	
Y	THOMAS ET AL: "Comparative receptor binding analyses of cannabinoid agonists and antagonists" J PHARMACOL. EXP THER, vol. 285, no. 1, 1998, pages 285-292, XP000942485 see whole document	1-10	
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Form PCT/ISA/210 (continuation of second sheet) (July 1992)

nternational application No.
PCT/EP 02/10434

INTERNATIONAL SEARCH REPORT

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of Irrst sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 1-3,9,10 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: see FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this international Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 1-3,9,10

Present claims 1-3,9,10 relate to a compound or use defined by reference to a desirable characteristic or property, namely that it be a prodrug of the formula I compounds.

The claims cover all compounds and uses having this characteristic or property, whereas the application provides support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT for only a very limited number of such compounds and uses. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Independent of the above reasoning, the claims also lack clarity (Article 6 PCT). An attempt is made to define the product by reference to a result to be achieved. Again, this lack of clarity in the present case is such as to render a meaningful search over the whole of the claimed scope impossible. Consequently, the search has been carried out for those parts of the claims which appear to be clear, supported and disclosed, namely those parts relating to the compounds of formula I and their use, and not their prodrugs.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

Information on patent family members

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